

CLAIMS

What is claimed is:

- 5 1. A method for assaying a sample for the presence of a target molecule comprising:
 providing a liquid sample suspected of comprising the target molecule;
 contacting the sample with a filter, said filter comprising a sensor molecule
 attached thereto, said sensor molecule capable of specifically binding to the target
 molecule, if present;
- 10 passing the sample transversely through said filter using a pressure-controlling
 apparatus under conditions that allow the sensor molecule to bind to the target molecule;
 recovering the remaining liquid sample; and
 determining whether the target has bound to the sensor.
- 15 2. The method of claim 1, wherein the sample is selected from the group consisting
 of blood; urine; semen; milk; sputum; mucus; pleural fluid; pelvic fluid; sinovial fluid;
 ascites fluid; a body cavity wash; eye brushing; skin scrapings; a buccal swab; a vaginal
 swab; a pap smear; a rectal swab; an aspirate; a needle biopsy; a section of tissue; plasma;
 serum; spinal fluid; lymph fluid; an external secretion of the skin, respiratory, intestinal,
20 or genitourinary tract; tears; saliva; a tumor; an organ; a microbial culture; and an *in vitro*
 cell culture constituent.
- 25 3. The method of claim 1, wherein the sensor comprises an antibody.
4. The method of claim 1, wherein the sensor comprises a polynucleotide.
5. The method of claim 1, wherein the sensor comprises a peptide nucleic acid.

6. The method of claim 1, wherein a plurality of different sensors are attached to the filter, wherein each of said plurality can selectively bind to a corresponding different target.

5 7. The method of claim 1, wherein the target is a cell surface molecule.

8. The method of claim 1, wherein the target is a soluble molecule.

9. The method of claim 1, wherein the target is membrane-bound.

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10. The method of claim 1, wherein the target is DNA.

11. The method of claim 1, wherein the target is RNA.

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12. The method of claim 1, wherein the target is from a pathological organism.

13.. The method of claim 1, wherein the target is a viral marker.

20 14. The method of claim 1, further comprising comparing a result from said determining to a result obtained from a control sample.

15. The method of claim 14, where the control sample is a positive control.

16. The method of claim 14, where the control sample is a negative control.

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17. The method of claim 1, further comprising washing said sample prior to said determining.

18. The method of claim 1, wherein the sample comprises a water-soluble alcohol in an amount effective to preserve the sterility of the solution toward at least one contaminant.
- 5 19. The method of claim 1, wherein determining whether the target has bound to the sensor comprises contacting the filter with a labeled secondary sensor, and determining whether label is associated with the filter.
- 10 20. The method of claim 19, wherein the first label comprises an agent selected from a chromophore, a lumiphore, a fluorophore, a chromogen, a hapten, an antigen, a radioactive isotope, a magnetic particle, a metal nanoparticle, an enzyme, an antibody or binding portion or equivalent thereof, an aptamer, and one member of a binding pair.
- 15 21. The method of claim 20, wherein the agent is an enzyme selected from alkaline phosphatase, horseradish peroxidase, β -galactosidase, glucose oxidase, a bacterial luciferase, an insect luciferase and sea pansy luciferase.
- 20 22. The method of claim 20, wherein the agent is a fluorophore.
23. The method of claim 22, wherein the fluorophore is a semiconductor nanocrystal.
24. The method of claim 23, wherein the fluorophore is a fluorescent dye.
25. The method of claim 20, wherein the agent is an enzyme, and a chemiluminescent substrate is used to detect the presence of agent.